The Nagoya Protocol International Access and Benefit Sharing Regime
By Viviana Munoz Tellez *

I. Introduction
This year marks the 10th year anniversary of the Nagoya Protocol on Access to Genetic Resources and the Fair and Equitable Sharing of Benefits Arising from their Utilization to the Convention on Biological Diversity. After six years of negotiations, in 2010 the elaboration of an international instrument on access and benefit-sharing was completed, concluding one of the most significant milestones of international environmental law in the last years. From October 2014, the Nagoya Protocol is in force.¹

The Convention on Biological Diversity (CBD) pursues the following three objectives: (i) the conservation of biological diversity, (ii) the sustainable use of its components and (iii) the fair and equitable sharing of the benefits arising out of the utilization of genetic resources (article 1 CBD). The Nagoya Protocol progresses in the implementation of the third objective.

The CBD advances the principle of national sovereignty of States over their natural resources. Such recognition resulted in a turning point regarding the legal nature of biological resources, which include “genetic resources, organisms or parts thereof, or any other biotic component of ecosystems with actual or potential use or value for humanity” (CBD Article 2). The CBD recognized for the

Abstract
The Nagoya Protocol on Access to Genetic Resources and the Fair and Equitable Sharing of Benefits Arising from their Utilization to the Convention on Biological Diversity entered into force in October 2014. Ten years have now passed since the adoption of the Protocol by the Parties to the Convention on Biological Diversity, now with 129 Parties. The Protocol requires countries to set up access and benefit sharing rules and procedures for the Protocol’s implementation at the national level. This policy brief describes the main characteristics of the Protocol and makes recommendations for countries to advance in its implementation. Importantly, the Protocol’s language empowers countries with considerable policy space for the design of domestic access and benefit-sharing rules.

* Programme Coordinator, Health, Intellectual Property and Biodiversity Programme (HIPB), South Centre. This Policy Brief updates South Centre Policy Brief No.18 (May 2015) by Viviana Munoz and Daniela Guarás.
first time that States have the authority to determine access to their genetic resources (CBD Article 15). Prior to the adoption of CBD, plant genetic resources were considered as a heritage of mankind being therefore freely accessible. This view is embodied in the FAO International Undertaking on Plant Genetic Resources (IUPGR) of 1983. Nevertheless, in the FAO context, it later recognized that the concept of mankind’s heritage, as applied in the IUPGR, is subject to the sovereignty of States over their plant genetic resources (FAO Resolution 3/91). This concept was also extended in the International Treaty on Plant Genetic Resources for Food and Agriculture (ITPGRFA). That said, the CBD and the ITPGRFA are distinct regimes, as the latter specifically applies to access and benefit sharing of plant genetic resources for food and agriculture.

In addition to legally empowering countries to control access to genetic resources, the CBD also defined two conditions to which access can be subject to, if so desired. These conditions are prior informed consent (PIC) and the establishment of mutually agreed terms (MAT). These conditions are meant to provide a basis for ensuring the third objective of the CBD, the fair and equitable benefit sharing arising out of the utilization of genetic resources. The inclusion of benefit sharing as an objective of the Convention responded to demands by developing countries. While developing countries were being asked to increase commitments to protect genetic resources, they sought to address a perceived historical imbalance whereby, for decades, they were mainly providers of genetic resources that were mainly exploited by and for the benefit of firms and other users in developed countries.

After CBD entered into force, the international community started building steps for the implementation of the first two objectives but not enough efforts were made to effectively implement the fair and equitable sharing of the benefits arising out of the utilization of genetic resources (ABS). This differentiation can be acknowledged through the various initiatives that encouraged countries to establish measurable targets for the conservation of biodiversity and sustainable use of its components, including genetic resources, as well as in the negotiations for the approval of the different Strategic Plans under CBD. However, as regards benefit sharing the main achievement was the approval of the non-binding Bonn guidelines on access to genetic resources and the fair and equitable sharing of the benefits arising from their utilization (Decision VI/24).

Developing countries were concerned about the misappropriation of their resources, usually known as ‘biopiracy’. Consequently, they demanded for an international regime on access and benefit sharing to be negotiated under the CBD in order to advance on the implementation of the Convention’s third objective while providing greater legal certainty for users and providers. In this context, countries started a negotiating process that resulted in the adoption and recent entry into force of the Nagoya Protocol.

The Protocol presents opportunities but also challenges. While rules on access and benefit sharing can and should be designed first and foremost at the national level, the CBD and the Protocol provide an important basis of international agreed rules that apply to all providers and users of genetic resources, in all countries that become a Party to the Protocol. Moreover, the flexibility in the language included in the Protocol provides countries with policy space for the design of domestic policies so as to maximise the benefits that arise from its implementation in accordance with local conditions. The vague language of the Protocol in many of its provisions reflects the extent to which compromises needed to be made in the negotiations to reach an agreed outcome. Hence, since the text may allow diverse interpretations, the interpretation of the Protocol should be carefully considered.

This policy brief describes the main characteristics of the Nagoya Protocol and highlights the core elements that developing countries need to bear in mind upon its ratification and subsequent implementation. Recommendations are made for countries to advance in its implementation.

II. What is the aim of the Nagoya Protocol?

The objective of the Protocol is the fair and equitable sharing of the benefits arising from the utilization of genetic resources (see Box 1). It applies to genetic resources under the scope of the CBD and the benefits arising from the utilization of those resources, as well as traditional knowledge (TK) associated with them.

The Nagoya Protocol expands on the obligations of the CBD on access and benefit-sharing (ABS), to effectively create an international ABS system. The Protocol is a detailed roadmap of international agreed principles and rules for the access and utilization of genetic resources and associated traditional knowledge. Through its implementation by governments, it is expected to provide users and providers of genetic resources and holders of traditional knowledge in all countries with greater clarity and certainty of what is permissible or not. Importantly, it clarifies the measures that countries can take to condition access to genetic resources and associated traditional knowledge, and requires commitments, including on tracking and monitoring of the utilization made of genetic resources and associated traditional knowledge, not only by countries that provide genetic resources but also by user countries, even when the latter choose not to regulate access to their genetic resources or associated traditional knowledge.

The fact that in some areas the agreed language of the Protocol is broad, such as in relation to the scope of the provisions of the Protocol covering derivatives, makes it imperative for countries to introduce an adequate interpretation in their domestic legislation. Fortunately, the Protocol allows countries sufficient policy space to define the details of their ABS laws at the national level.
III. Main obligations for Parties to implement the Protocol

The Protocol clarifies rights and obligations for both providers and users related to the access to genetic resources and traditional knowledge associated with those resources as well as for benefit-sharing and monitoring of the utilization of genetic resources. Some of the main elements are included below.

III.1. Genetic resources

III.1.a. Access

Based on the principle of sovereignty of States over their natural resources, the fundamental provision of the Protocol with regards to access to genetic resources and traditional knowledge associated with those resources as well as for benefit-sharing and monitoring of the utilization of genetic resources. Some of the main elements are included below.

Box 1. What is benefit-sharing?

The notion of benefit-sharing finds its roots in the third objective of the CBD that is the “fair and equitable sharing of the benefits arising out of the utilization of genetic resources, including by appropriate access to genetic resources and by appropriate transfer of relevant technologies”. The CBD in Article 15.7 affirms that each Party must take measures with the aim of sharing, upon mutually agreed terms, in a fair and equitable way the results of research and development and the benefits arising from the commercial and other utilization of genetic resources with the country providing such resources. This is at the core of the Nagoya Protocol.

The goal of benefit-sharing is that users of genetic resources effectively share monetary and non-monetary benefits derived from the access and utilization of those resources with the country providing those resources (countries of origin, see box 2). As regards traditional knowledge associated to those resources, benefits should be shared with the communities that are holders of that knowledge in accordance to the measures that need to be taken to this end at the national level (article 5.2 of the Protocol).

Due to details that the Protocol contains compared to provisions included in the CBD, it is an important guidance for countries that need to design their national ABS legislations. Furthermore, it gives countries the possibility of prioritising ABS related issues in their national agendas.

On the other hand, countries that already have ABS legislation would need to evaluate to what extent they will have to adapt their existing legislation to the requirements of the Protocol. Moreover, they will need to discuss what the best approach could be in order to use the policy space provided by the Protocol and, subsequently, decide whether to ratify or not.

The Nagoya Protocol includes additional definitions to those found in the CBD, including for the terms “utilization of genetic resources” and “derivatives”. Importantly, the inclusion of such definitions clarifies that the Nagoya Protocol includes under the scope of its obligations utilization of genetic resources and their derivatives. The language of “utilization of genetic resources” refers specifically to the conduct of research and development, including their “derivatives”, that is, naturally occurring biochemical compounds even when they do not contain functional units of heredity. Activities that fall outside this scope, such as trade in commodities, are not covered. For legal certainty, it will be important that developing countries introduce these definitions in their national ABS legislation.

Box 2. Some relevant definitions

The definitions of the CDB (included in article 2 of the CDB) apply to the Nagoya Protocol. In addition, some terms such as “utilization of genetic resources” and “derivatives” are defined in the Protocol.

- **Biotechnology**: “any technological application that uses biological systems, living organisms, or derivatives thereof, to make or modify products or processes for specific use”
- **Country of origin of genetic resources**: “the country which possesses those genetic resources in in-situ conditions”
- **Country providing genetic resources**: “the country supplying genetic resources collected from in-situ sources, including populations of both wild and domesticated species, or taken from ex-situ sources, which may or may not have originated in that country”
- **Derivatives**: “a naturally occurring biochemical compound resulting from the genetic expression or metabolism of biological or genetic resources, even if it does not contain functional units of heredity”
- **Genetic material**: “any material of plant, animal, microbial or other origin containing functional units of heredity”
- **Genetic resources**: “genetic material of actual or potential value”
- **Utilization of genetic resources**: “to conduct research and development on the genetic and/or biochemical composition of genetic resources, including through the application of biotechnology as defined in Article 2 of the Convention”
The Protocol places responsibility on both provider and user countries to take measures so to ensure that prior informed consent has been obtained previous to the access to genetic resources and/or the associated traditional knowledge, as well as to guarantee the involvement of indigenous and local communities in that process, when relevant.

Prior informed consent means that a user (researcher, firm, etc.) that seeks access to a genetic resource or traditional knowledge associated to the resource needs to receive express acceptance or permission from the country providing genetic resources (whether or not it is the country of origin of the genetic resource), or an indigenous or local community providing traditional knowledge associated to those resources, as may be the case according to national legislation. The consent is materialised through the issuance of a permit for that access. According to the Protocol, each country can decide whether to regulate access to its genetic resources and how to do it.

In particular, countries providing genetic resources that require prior informed consent have some obligations to comply with, as follows:

- Domestic ABS legislation, rules and procedures need to provide legal certainty, clarity and transparency, be fair and non-arbitrary
- Information on how to apply for prior informed consent need to be provided as well as a clear and transparent written decision by a competent national authority (in a cost-effective manner and within a reasonable period of time)
- A permit has to be issued at the time of access, as evidence of the decision to grant prior informed consent and of the establishment of mutually agreed terms (MAT). This will also be notified to the ABS Clearing-House.iii

These obligations on providers are meant to facilitate the awareness of the conditions/requirements that users need to comply with prior to accessing genetic resources in each provider country, which might vary from one to the other.

Prior informed consent is directly connected to the establishment of mutually agreed terms (MAT) for benefit sharing in relation to the utilization of genetic resources and/or associated traditional knowledge. MAT means that the conditions for the access and utilization of the resources and/or traditional knowledge, such as the benefits to be shared from the utilization, have been fairly and equitably negotiated and agreed among the Party providing the resource or traditional knowledge holder and the user. Provider countries must establish clear rules and procedures for the establishment of MAT. These terms refer to the conditions of the ‘contract’ between provider and user, and may include but are not limited to monetary and other forms of benefit-sharing.

While generally access can be conditioned to PIC and MAT, the Protocol includes a reference to some instances that can benefit from facilitated access (article 8). These situations include creating conditions to promote research, including through simplified measures on access for non-commercial research purposes as well as paying due regard to cases of present or imminent emergencies that threaten or damage human, animal or plant health, as determined nationally or internationally. It also mandates Parties to consider the importance of genetic resources for food and agriculture in implementing its access and benefit-sharing legislation or regulatory requirements.

III.1.b. Fair and equitable benefit-sharing

The Protocol’s language on fair and equitable benefit-sharing is based on CBD’s provisions on this matter. Nevertheless, it goes beyond the CBD in two specific points. Firstly, as it provides the opportunity that not only benefits arising from the utilization of genetic resources but also those from ‘subsequent applications and commercialization’ are shared with the provider country (Article 5, see box 4). Secondly, it recognises that in some jurisdictions, genetic resources can be held by indigenous and local communities. In these situations, countries need to establish means to ensure that benefits arising from the utilization of those genetic resources are shared with the communities in a fair and equitable manner. Benefits to be shared are both monetary and non-monetary. A non-exhaustive list of possible benefits is included in the Annex of the Protocol. Conditions and mechanisms for benefit-sharing will be based on MAT between provider and user. Thus, the enhancement of institutional and human capabilities to negotiate beneficial conditions is a central issue to be addressed domestically.

III.2. Traditional Knowledge associated with genetic resources

III.2.a. Access and benefit sharing

Access to and utilization of traditional knowledge associated with genetic resources is also a cross-cutting issue throughout the text. Considering the characteristics of traditional knowledge, i.e. the importance of indigenous and local communities as holders of that knowledge, PIC but the establishment of MAT acquire fundamental importance.

Although provisions on TK associated to genetic resources, in the Protocol are not as detailed as for the case of genetic resources, the Protocol substantially improves upon the CBD language. Article 7 of the Protocol, in complementing Article 8(j) of the CBD, requires that Parties take measures, as appropriate, to ensure that TK associated with genetic resources that is held by indigenous and local communities is accessed with PIC or their approval.
and involvement, and that MAT have been established. This applies to both provider and user countries.

The notion of benefit-sharing related to the utilization of associated traditional knowledge (TK) is another element in which the Protocol reinforces CBD’s provisions on TK, by reference to the establishment MAT, following PIC. Moreover, while the CBD only expresses the desire for benefit-sharing to exist for the utilization of TK, the Protocol supersedes that provision and sets a concrete obligation.

Article 12 of the Protocol requires consideration to indigenous and local communities’ customary laws, community protocols and procedures with regard to associated TK. It is a big step in which the Protocol advances CBD. However, since the article provides some flexibility, each country needs to decide in accordance with its national legislation whether to implement it and how to do it.

It can be challenging but nonetheless it is of great importance for governments to establish effective mechanisms to ensure that PIC and MAT for benefit-sharing are established among providers and users prior to the access, and utilization of TK associated with genetic resources. Challenges include the fact that the traditional knowledge associated to the genetic resource may be found outside the community, e.g. libraries, repositories, and the geographical location of indigenous or local communities, when they are located in remote areas with no easy access and appropriate communication technologies available. In this regard, it is crucial for governments to work together with indigenous and local communities to find viable options to develop a workable system.

The Nagoya Protocol makes no distinction between TK that is well-known outside the indigenous or local community, from TK that is undisclosed or held secret. Thus, it can be understood that PIC and MAT is required for access to any TK associated to GR to be lawful, though countries can define the matter in national legislation.

III.3. Monitoring and Compliance

One of the most important aspects of the Protocol is that it introduces the obligations on monitoring and compliance. It is the first time that an international instrument includes international rules on monitoring and compliance in user countries on access to and utilization of genetic resources and associated TK.\(^v\)

Article 15 and 16 refers to compliance with domestic ABS legislation or requirements regarding access to genetic resources and associated TK respectively. User and provider countries alike have to introduce measures for ensuring that genetic resources or associated traditional knowledge utilized in their jurisdiction have been accessed in accordance with PIC and that MAT have been established, as required by domestic ABS legislation or regulatory requirements of the country providing genetic resources or associated TK, or where indigenous and local communities providing associated TK are located (Articles 15, 16).\(^vi\)

Importantly, the Protocol requires that associated TK is accessed in accordance with PIC or with the approval and involvement of indigenous and local communities.

The Protocol also mandates Parties to take measures to address cases of non-compliance with domestic ABS legislation, and that importance of establishing cooperative procedures and institutional mechanisms to promote compliance with the provisions of the Protocol and to ad-
The Nagoya Protocol International Access and Benefit Sharing Regime

dress cases of non-compliance.

Parties are also required to monitor the utilization of genetic resources in their territories, as a means to support compliance, and to enhance transparency on the utilization that is being made of genetic resources (Article 17). In this regard, the Protocol obliges countries to establish at least one ‘checkpoint’ that would be able to collect or receive information on prior informed consent, the source of the genetic resources or the establishment of mutually agreed terms. Nonetheless, no list of possible checkpoints is included in the Protocol. To complete this task, countries providing genetic resources can ask for users to provide this information to the national authorities.

In view of the lack of any indicative list in the Protocol, there is flexibility to designate the checkpoint/s that each country considers more appropriate for the effective completion of the tasks. The only indication provided by the Protocol is that checkpoints should be relevant to the utilization of genetic resources, or to the collection of relevant information at any stage of research, development, innovation, pre-commercialisation or commercialisation.

An option that was discussed during the negotiations prior to the adoption of the Protocol was to explicitly mention patent offices, market approval authorities or research funding institutions as checkpoints. Although no agreement was reached for that inclusion, all options are now available for countries' consideration when implementing the Protocol domestically. For instance, in some countries the responsibility is shared between patent offices and environmental agencies. Likewise, considering the characteristics of each country, in those cases in which genetic resources are regulated by the local governments, additional layers of responsibility exist.

Article 17 only refers to utilization of genetic resources, making no reference to monitor the utilization of associated TK. In this regard, additional international instruments could be considered as a complement of the Protocol’s provision. One of those complementary regimes can be the elaboration of a(n) international legal instrument(s) to ensure the effective protection of TK, traditional cultural expressions (TCEs) and genetic resources that has been taking place in WIPO.

Compliance and monitoring are closely related issues because effective compliance requires monitoring. It is of vital importance that both user and provider countries increase their capacity to monitor the utilization of both genetic resources and associated traditional knowledge to effectively implement the ABS system. However, it must also be recognized as a challenge that countries face, in particular developing countries, i.e. in cases when the genetic resources were accessed ex situ, the associated TK is not directly attributable to a single indigenous or local community, or in the case of genetic resources shared across-borders, it is difficult to establish when and from where the access to the genetic resource was made.

The utilization of genetic resources includes a wide variety of sectors such as pharmaceuticals, food, agriculture, biotechnology or cosmetics and existing differences between them can, and should, be addressed at the domestic level. For example, in sectors such as agriculture it is sometimes difficult to determine the origin of genetic resources. In particular, considering the relevance that the paradigm of plant genetic resources as a common heritage had for that sector and that exchanges of materials throughout the decades have been multiple, the identification of a country of origin can be something challenging. Furthermore, it is frequent that genetic materials utilized for food and agriculture are stored and conserved ex situ, in banks such as CGIAR international collections.

To some extent, it can be interpreted that this kind of situations would fall under the global multilateral benefit-sharing mechanism that is under development. However, article 10 only refers to the utilization of genetic resources and associated TK that occur in transboundary situations or for which it is not possible to grant or obtain prior informed consent.

Different dynamics characterise the way in which diverse sectors work. But what does it mean in terms of ABS? Some examples of the characteristics of private activities of two sectors that can be subject to comply with ABS legislation are mentioned below.

- Pharmaceutical sector: Some changes are being acknowledged with regard to the way the pharmaceutical sector conducts research and development. In the last years, there has been an increase in the demand microorganisms moving away from the traditional demand for plants.
- Agricultural sector: It is one of the sectors that continue being more dependent on access to genetic resources. Again, the smaller companies are those that require access to public collections since the large ones usually have their own collections of plant genetic resources. However, there is an additional aspect that can affect or regulate the exchange of materials in this sector. In this regard, it is relevant to highlight that the Nagoya Protocol recognises in its Preamble the role of the International Treaty on Plant Genetic Resources for Food and Agriculture (ITPGRFA) of the Food and Agriculture Organization and its multilateral ABS system. In addition, article 4 states that the Protocol shall not apply for the specific genetic resources covered by a specialised international ABS instrument. This means that plant genetic resources for food and agriculture under the scope of the ITPGRFA may be exempted from the establishment of MAT under the Protocol. Therefore, national legislation frameworks should clarify the scope of both agreements so to create a mutually supportive relationship between them.
IV. Intellectual Property aspects in the Nagoya Protocol

The tensions and links that exist between genetic resources, technological capacities and intellectual property have long been recognized in international debates, i.e. Brundtland Report (1987). Whereas the linkages between biodiversity and intellectual property have been expressly recognised in the CBD and the FAO ITPGRFA, processes are underway in other international organisations such as the World Trade Organization (WTO) and World Intellectual Property Organization (WIPO) on the relationship between these two areas. However, no decisions have been adopted so far, i.e. in the WTO discussions on how to create a mutually supportive relationship between the CBD and the Agreement on Trade-Related Aspects of Intellectual Property Rights (TRIPS) and the protection of traditional knowledge, and in the WIPO on how to create an effective system of protection for genetic resources, traditional knowledge and traditional cultural expressions.

The controversies around intellectual property (IP) arise due to tensions between the interests of providers; actors that have legal rights to control access to genetic resources and associated traditional knowledge, i.e. indigenous communities, States’ institutions, and the interests of users whose activities include research, development and commercialization of genetic resources, including their derivatives. Such tensions arise when genetic resources or associated TK are granted IPR protection. IP rights (IPRs), on the one hand, can serve as a tool to generate economic benefits for the owner of the IPR-protected goods. On the other hand, IPRs exclude third parties from the unauthorized use of the protected good for as long as the period of protection of the IPR lasts, even when a third party may have contributed to the conservation of, or provided important knowledge for the subsequent utilization of the genetic resources.

During the negotiations of the Nagoya Protocol, many developing countries pushed for concrete language on IP. Proposals included to recognize the role of patent offices acting as checkpoints in order to assist in ensuring compliance with national ABS laws, and in particular to introduce a disclosure requirement in patent and other IPRs applications so to make explicit the country of origin of genetic resources and associated TK and the information contained in the MAT. Other proposals included the use of patent databases to track both the utilization of genetic resources as well as the benefit-sharing obligations declared in the MAT. Although measures concerning IPRs are not directly mentioned in the final text of the Protocol, these can be included as part of the implementation of the Protocol where relevant, for example as part of measures for monitoring and compliance.

It is thus important that in national legislation, countries introduce specific language to designate IP offices as a checkpoint (among other checkpoints, i.e. customs offices) for the purposes of monitoring and compliance with ABS legislation, consistent with Article 17 of Nagoya. The function of checkpoints would be to receive information concerning the utilization of genetic resources or their derivatives. This would be done through the mandatory requirement to disclose in a patent or other IPR applications, i.e. plant variety protection applications, the source of the genetic resources or derivatives. To date, various countries have established IP offices as checkpoints.

V. Other obligations in the Nagoya Protocol

The Nagoya Protocol requires that certain institutional arrangements are put in place, particularly to facilitate transparency. Every Party has to designate a national focal point as well as a competent national authority/ies for activities that range from communication with the Secretariat of the Protocol to the implementation of the Protocol itself (Article 13). Additionally, it is necessary to make available relevant information for the ABS clearing-house mechanism. This includes legislative, administrative and policy measures on ABS, information on the national focal point and competent national authority/ies and the permits issued at the time of access as evidence of the decision to grant PIC and of the establishment of MAT. xii

It is worth noting that while the Protocol leaves many areas to be developed at the domestic level, various arrangements still need to take place in the international arena to provide the necessary basis for an ease implementation. The evolvement of these aspects will take

---

Box 5. IP offices as checkpoints

Some examples of countries that have notified to the ABS Clearing-House Mechanisms that they have established IP offices as one instance of checkpoint for compliance with national ABS laws:

- Bhutan: Department of Intellectual Property, Ministry of Economic Affairs.
- Ecuador: Servicio Nacional de Derechos Intelectuales (Senadi).
- Peru: Dirección de Invenciones y Nuevas Tecnologías (DIN) del Instituto Nacional de Defensa de la Competencia y de la Protección de la Propiedad Intelectual (INDECOPI).  
- Switzerland: Swiss Federal Institute of Intellectual Property  
- Uruguay: Dirección Nacional de la Propiedad Industrial del Ministerio Industria Energía y Minería.
place through the decisions made by the Conference of the Parties serving as the Meeting of the Parties to the Nagoya Protocol (COP-MOP).

VI. Outstanding issues under discussion by the Parties to the Protocol

There are several important issues that are currently being discussed among the Conference of the Parties (NP-MOP). One is the treatment of digital sequence information on genetic resources for purposes of ABS, given the increasing generation and use of digital sequence information on genetic resources and its publication in both public and private databases, advances in data analytics and new technologies for the current and future utilization of genetic resources. Many countries have adopted domestic measures that regulate the access to and use of digital sequence information on genetic resources as part of their access and benefit-sharing frameworks, but Parties are yet to come to agreement on how to address this question. A working group is now examining the matter and will report back to the NP-MOP in 2021.\textsuperscript{viii}

Another issue under consideration by the NP-MOP is the establishment a global multilateral benefit-sharing mechanism cases of genetic resources and traditional knowledge associated with genetic resources that occur in transboundary situations or for which it is not possible to grant or obtain prior informed consent, in accordance to Article 10.\textsuperscript{xiv}

The NP-MOP is also considering criteria for identifying specialized international access and benefit-sharing instruments in the context of Article 4, paragraph 4, of the Nagoya Protocol. The potential criteria are under discussion and have not been agreed by Parties to the Protocol.\textsuperscript{vi} Currently Parties are exchanging views.\textsuperscript{vii} An important consideration is whether for sectors and situations, such as health emergency, where expedited access is desirable, a global multilateral benefit-sharing mechanism should be established.

The Fourth meeting of the Conference of the Parties serving as the meeting of the Parties to the Nagoya Protocol on Access to Genetic Resources and the Fair and Equitable Sharing of Benefits Arising from their Utilization (COP-MOP 4) is expected to meet in the second half of 2021 in Kunming, China.

VII. Assessment and review of the effectiveness of the Nagoya Protocol

A first assessment and review of the effectiveness of the Nagoya Protocol was undertaken by the NP-MOP in 2018, in accordance to Article 31.\textsuperscript{xvii} The review notes the progress achieved yet points to the gaps by Parties in establishing the necessary access and benefit-sharing (ABS) legislative, administrative and policy measures and institutional arrangements. A particular challenge identified is the implementation of some of the new elements of the Protocol, namely the provisions on compliance, monitoring the utilization of genetic resources, including the designation of checkpoints, and the obligations related to indigenous peoples and local communities. Many countries are yet to designate checkpoints, which is essential for effective monitoring and compliance with domestic ABS regulations. Many Parties have also not published information to the ABS Clearing-House.

The review further suggests that Parties need to increase efforts for the participation of indigenous peoples and local communities and relevant stakeholders (e.g. different business sectors and the scientific community) as well as coordination among different institutions and ministries (e.g. science and education, agriculture, trade, intellectual property).

The review also finds that many Parties still lack the necessary capacity and financial resources to make the Protocol operational. While there are several capacity-building and development initiatives, these are insufficient and thus more capacity-building and financing support is necessary.

The second assessment and review of the effectiveness of the Protocol will be held in 2024.

VIII. Policy Recommendations

The Nagoya Protocol is an important international instrument on ABS under the CBD. The Protocol was concluded 10 years ago and has been in force for 6 years. However, many of the provisions of the Protocol require implementation in national legislation. Thus, countries need to establish and/or revise laws and regulations (and related laws), institutional structures, administrative and policy measures on ABS. The implementation of the Protocol through national measures by all Parties, and their transparent reporting, is essential to materialise the expected benefits of an international regime on ABS that creates more certainty for both providers and users of genetic resources and associated traditional knowledge. The obligations in the Protocol apply to all Parties, whether they are countries that are mainly users or providers of genetic resources.

To conclude, some elements are highlighted and recommendations provided to encourage discussions at the national level on what the most appropriate ways to implement the Protocol might be:

- The Protocol does not replace the need of countries to develop legislation on ABS.\textsuperscript{xviii} It is rather a basis for its development. Attention needs to be given to the interpretation of the Protocol provisions so to maximise the policy space provided by them, and in the measures adopted for implementation. A careful review of the countries’ interests of both providers and users of genetic resources and associated traditional knowledge is necessary. Sharing of experiences among countries in implementing the Nagoya Protocol, in particular South-South cooperation, is important to support development of national ABS regimes.
- Dynamics that characterise the exchange of mate-
rials within each sector have to be understood. Some studies indicate that whereas large companies access to materials in the past and in some cases have their own collections of plant genetic resources, different trends may characterise other resources such as marine and terrestrial microorganisms. This reality, while increasing the difficulties to monitor the utilization of genetic resources and associated TK, also highlights the importance of being conscious about who the targeted actors are in each case so to design the most appropriate and effective measures.

- Frequently, many governmental agencies are directly or indirectly involved in the implementation of genetic resources related policies. It is thus imperative that different ministries and departments coordinate and cooperate between them for the successful implementation of ABS measures.

- It is important for countries providing genetic resources and associated TK to raise awareness about their domestic ABS rules so that all potential providers, i.e. indigenous and local communities, gene banks, as well as users are aware of the conditions that they must comply with. Developing countries have to make use of the provisions of the Protocol in this matter.

- Capacity-building is central to create an appropriate institutional base for the implementation of the agreement. In particular, to empower providers to negotiate with potential users for the access based on PIC and for the establishment of MAT for benefit-sharing.

- Developing countries should actively participate in the continued deliberations taking place at the international level with regards to the implementation of the Nagoya Protocol, provisions where follow-up by the COP MOP is necessary, and related processes in other international fora, particularly the WTO, WIPO and FAO. In particular, experiences of the implementation at the national level should be shared so to facilitate a smooth implementation of the Protocol following its entry into force.

Endnotes:

1 For more information on the status of accessions/ratifications to the Nagoya Protocol, see [https://www.cbd.int/abs/nagoya-protocol/signatories/](https://www.cbd.int/abs/nagoya-protocol/signatories/).

2 The “country providing genetic resources” and the “country of origin” are two distinct concepts. Both terms are defined in the CBD and those definitions apply for the Nagoya Protocol (See Box 2).


5 Although the Protocol refers to Parties in generic terms, the content of the provisions make user countries as important actors for the implementation of articles 15 and 16.


9 Ibidem at viii.


12 See Decision 3/12 of the NP - MOP 3, 2018, at: [https://www.cbd.int/decision/np-mop/?id=13700](https://www.cbd.int/decision/np-mop/?id=13700).

13 See Decision 3/10 of the NP - MOP 3, 2018 at: [https://www.cbd.int/decision/np-mop/?id=13695](https://www.cbd.int/decision/np-mop/?id=13695).


15 See for example, the submission by the African Union, [https://www.cbd.int/abs/submissions/Art4-4/2019/AfricanUnion.pdf](https://www.cbd.int/abs/submissions/Art4-4/2019/AfricanUnion.pdf).


18 Ibidem at viii.

The Nagoya Protocol International Access and Benefit Sharing Regime

<table>
<thead>
<tr>
<th>Previous South Centre Policy Briefs</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. 78, May 2020 — The 73rd World Health Assembly and Resolution on COVID-19: Quest of Global Solidarity for Equitable Access to Health Products by Nirmalya Syam, Mirza Alas and Vitor Ido</td>
</tr>
<tr>
<td>No. 79, June 2020 — Articles 7 and 8 as the basis for interpretation of the TRIPS Agreement by Thamara Romero</td>
</tr>
<tr>
<td>No. 80, June 2020 — Intellectual Property, Innovation and Access to Health Products for COVID-19: A Review of Measures Taken by Different Countries by Nirmalya Syam</td>
</tr>
<tr>
<td>No. 81, July 2020 — The UN General Assembly Resolutions on COVID-19: Solemn Assurances for Access to Health Technologies without an Action Plan by Nirmalya Syam</td>
</tr>
<tr>
<td>No. 82, July 2020 — Examining antimicrobial resistance in the light of the COVID-19 pandemic by Mirfin Mpundu, Caline Mattar and Mirza Alas</td>
</tr>
<tr>
<td>No. 83, August 2020 — United States: An Obsolete Trade Practice Undermines Access to the Most Expensive Drugs at More Affordable Prices by Maria Fabiana Jorge</td>
</tr>
<tr>
<td>No. 84, September 2020 — A New Trend in Trade Agreements: Ensuring Access to Cancer Drugs by Maria Fabiana Jorge</td>
</tr>
<tr>
<td>No. 85, October 2020 — Política de industrialización de litio, el caso boliviano por Hortensia Jimenez Rivera</td>
</tr>
</tbody>
</table>

The South Centre is the intergovernmental organization of developing countries that helps developing countries to combine their efforts and expertise to promote their common interests in the international arena. The South Centre was established by an Intergovernmental Agreement which came into force on 31 July 1995. Its headquarters is in Geneva, Switzerland.

Readers may reproduce the contents of this policy brief for their own use, but are requested to grant due acknowledgement to the South Centre. The views contained in this brief are attributable to the author/s and do not represent the institutional views of the South Centre or its Member States. Any mistake or omission in this study is the sole responsibility of the author/s. For comments on this publication, please contact:

The South Centre
International Environment House 2
Chemin de Balexert 7-9
PO Box 228, 1211 Geneva 19
Switzerland
Telephone: (4122) 791 8050
south@southcentre.int
https://www.southcentre.int

Follow the South Centre’s Twitter: South_Centre